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#### (57) Abstract

A process for altering the target insect range (spectrum) of pesticidal toxins which comprises recombining in vivo the hypervariable regions of two genes encoding a pesticidal toxin and having enough residual homology as to be able to promote in vivo recombination. According to the present invention, truncated genes obtained from well known strains of Bacillus thuringiensis variety kurstaki and separated by an antibiotic resistant marker gene - or part of it - are cloned in a plasmid vector which is then introduced in a strain of E. coli. In vivo recombination between the hypervariable regions of the toxins genes reconstitutes an entire hybrid toxin gene. Polypeptides encoded by these new hybrid toxin genes have different biological activity and an altered target insect range as compared to their parental toxin.

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insensitive.

NEW FUNCTIONAL BACILLUS THURINGIENSIS HYBRID GENES OBTAINED BY IN VIVO RECOMBINATION

- Bacillus thuringiensis is a spore-forming bacterium which, upon sporulation, produces an insecticidal parasporal crystal. Many <u>Bacillus thuringiensis</u> subspecies have been isolated and the vast majority of the strains which have been tested have shown specific activity only against larvae of Lepidopteran insects such as <u>Manduca sexta</u>, <u>Heliotis virescens</u> and <u>Trichoplusia ni</u>. Of the Lepidopterans, not all are equally sensitive to <u>Bacillus thuringiensis</u> For example, Spodoptera species tend to be relatively
  - The Lepidopteran specific <u>Bacillus thuringiensis</u> strains have been categorized according to flagellar serotype, crystal serotype as well as activity spectrum against various insects (Dulmage, 1981).
- 20 Among the better studied varieties of Lepidopteran specific <u>Bacillus thuringiensis</u> are <u>B.t. kurstaki</u> HD1 which is the strain used in "Dipel", B.t. HD 73, <u>B.t. dendrolimus</u>, <u>B.t. sotto</u>, <u>B.t. Berliner</u>.
- In recent years, new types of <u>Bacillus thuringiensis</u>

  25 with novel insecticidal specificities have been discovered. <u>B.t. israelensis</u> is toxic to larvae of several Dipteran species (mosquitoes and black flies), but not to Lepidopteran larvae (Goldberg and Margaht, 1977). More recently, two Coleopteran specific strains, <u>B.t. tenebrionis</u> and <u>B.t. San Diego</u>, which later were shown to be the same strain

### SUBSTITUTE SHEET

(Krieg et al., 1987), have been described (Krieg et

al., 1983, Krieg et al., 1984; Herrnstadt et al., These strains have shown activity against Colorado potato beetle and other Coleopteran pests.

entomopathogenic activity of the 5 thuringiensis parasporal crystal is due to its composition: in the case of the Bacillus thuringiensis subspecies specifically active against Lepidopteran larvae, it is composed of 130 to 160 Kdal protoxin polypeptides. Different subspecies and

The

- often individual strains of the same subspecies 10 produce endotoxins having a characteristic spectrum insect toxicity (Whiteley and Schnepf, 1986; of Andrews et al., 1987).
- For many years Bacillus thuringiensis has served as the basis of successful biological insecticides. 15 produce these insecticides, Bacillus thuringiensis is fermented until spores and crystals are obtained. The mixture of spores and crystals is then formulated to allow effective application on crop plants.
- Current, traditional <u>Bacillus</u> thuringiensis products 20 are in fact an example of classical industrial microbiology. These products are created through such traditional microbiological practices as strain isolation and improvement, and fermentation 25 optimization.
  - Two features of Bacillus thuringiensis have made it a popular and useful insecticide. First. Bacillus thuringiensis is considered extremely safe: it is harmless to humans, animals and useful insects.
- Second, Bacillus thuringiensis is a highly specific 30 insecticide: most strains of Bacillus thuringiensis show toxicity to only a single order of insects

(Lepidoptera or Coleoptera or Diptera), as indicated above.

These features have also made <u>Bacillus</u> thuringiensis an attractive target for biotechnology and recently the new tools of this modern technique, such as gene cloning and DNA sequencing, have begun to be applied to <u>Bacillus</u> thuringiensis, thus creating novel <u>Bacillus</u> thuringiensis pesticidal proteins with either more specificity, more toxic activity or an

- 10 altered range of toxicity for the host insect.

  The insecticidal activity of <u>Bacillus thuringiensis</u> resides in the parasporal crystal (Angus, 1954).

  Intact crystals can be isolated from sporulated cultures of <u>Bacillus thuringiensis</u> by density

  15 gradient centrifugation and these isolated crystals
- of the Lepidopteran specific <u>Bacillus thuringiensis</u> variety <u>kurstaki</u> were shown to be composed by protein subunits of approximately 130,000 daltons (Bulla et al. 1977). In some strains such as <u>Bacillus</u>
- thuringiensis variety <u>kurstaki</u> HD-73 there appears to be a single protein subunit, while in other strains such as <u>Bacillus thuringiensis</u> HD1 there appear to be two or more very similar proteins in the crystal (Wilcox et al., 1986).
- 25 The 130.000 dalton protein is considered to be a protoxin because it is toxic to larvae only after injection, but not after injection. The protoxin can be converted to the active toxin by digestion with proteases; and it has also been possible to isolate
- 30 proteolytic fragments of the protoxin which retain full toxic activity.

Bulla et al., 1981, found that a 68,000 dalton toxin

fragment could be derived from <u>Bacillus</u> <u>thuringiensis</u> <u>kurstaki</u> crystals upon prolonged incubation of the solubilized protein.

Crystals of the Coleopteran specific <u>Bacillus</u>

<u>5 thuringiensis</u> strains also appear to be composed of a single protein subunit, but of a much smaller size than the Lepidopteran protoxin.

Bernhard, 1986, isolated a 68,000 dalton protein from crystals of <u>Bacillus thuringiensis</u> <u>tenebrionis</u>, and

- 10 Herrnstadt et al., 1986, observed a 64,000 dalton protein from <u>Bacillus thuringiensis San Diego</u> crystals. These isolated proteins are toxic upon ingestion by sensitive Coleopterans.
- The crystal protein of the Lepidopteran specific

  Bacillus thuringiensis var. kurstaki strain, produced during the sporulation period, is also known as endotoxin, and around 20-30% of the cell protein synthetizing activity during sporulation is devoted to the production of this toxin.
- 20 Much work has recently been directed to the isolation and characterization of genes encoding <u>Bacillus</u> thuringiensis toxins. The analysis of such cloned genes has already yielded important insights into toxin structure and function: it has been shown for example that genes for the crystal proteins are located on large plasmid in addition to chromosomal DNA.

Several groups have reported cloning genes for Lepidopteran specific toxins. Most of these genes 30 have been cloned in <u>E. coli</u>, either utilizing antibodies to purified toxin to detect expression of the toxin, or utilizing synthetic oligonucleotide

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probes based on the toxin aminoacid sequence to detect toxin genes by hybridation. The genes cloned include several genes from <a href="Bacillus thuringiensis">Bacillus thuringiensis</a> kurstaki HD1 (Schnepf and Whiteley, 1981; Held et al., 1982; Watrud et al., 1985; Shivakumar et al., 1986; Thorne et al., 1986), and genes from <a href="Bacillus thuringiensis kurstaki">Bacillus thuringiensis kurstaki</a> HD 73 (Adang et al., 1985), from <a href="Bacillus thuringiensis sotto">Bacillus thuringiensis sotto</a> (Shibano et al., 1985), <a href="Bacillus thuringiensis Berliner">Bacillus thuringiensis Berliner</a> (Klier et al.,

- 10 1982; Wabiko et al., 1986), <u>Bacillus thuringiensis</u>
  <u>aizawa</u> (Klier et al., 1985) and <u>Bacillus</u>
  <u>thuringiensis thuringiensis</u> (Honigman et al., 1986).

  In general these genes have been shown to express
  toxin in <u>E. coli</u> and extracts of <u>E. coli</u> harbouring

  15 these genes are toxic to Lepidopteran larvae.
- The cloned toxin genes have been used as molecular probes to determine the toxin gene number and type of many Lepidopteran active <u>Bacillus thuringiensis</u> strains (Kronstad et al., 1983). This analysis has 20 shown that, while some strains (e.g. <u>Bacillus thuringiensis kurstaki HD73</u>) contain only a single toxin gene, many other strains contain multiple genes. <u>Bacillus thuringiensis kurstaki HD-1</u> (the Dipel strain) has three distinct toxin genes (Wilcox et al., 1986).

DNA sequences and derived aminoacid sequences of the toxin proteins have been determined for several of these genes. All the genes encode proteins of between 1156 and 1178 aminoacids which are largely homologous. In some cases genes isolated from strains which had been considered distinct varieties have been found to be nearly identical. For example, the

Bacillus thuringiensis kurstaki HD-1 gene sequenced by Schnepf et al (1985) is nearly identical to the Bacillus thuringiensis sotto gene of Shibano et al. (1985). Similarly, a Bacillus thuringiensis Berliner gene (Wabiko et al., 1986) is nearly identical in sequence to another Bacillus thuringiensis kurstaki HD-1 gene. On the other hand, a third gene from Bacillus thuringiensis kurstaki HD-1 (Thorne et al. 1986) is clearly different in sequence from the two mentioned above.

Cloning and sequencing of the structural genes the protoxin production from distinct strains Bacillus thuringiensis kurstaki have, therefore, revealed that different related genes are responsible for the synthesis of the crystal protein toxin. These 15 differences are evident not only among genes from different strains, but also among the multiple copies of the protoxin gene in the same strain (Andrews et 1987). The <u>kurstaki</u> HD-1 Dipel protoxin gene (Schnepf et al. 1985) and the <u>kurstaki</u> HD-73 protoxin 20 gene (Adang et al., 1985) show an homology of 85% at the primary DNA sequence level. K-1 type and K-73 type crystals show distinct toxic activity against different insect species (Jaquet et al., 1987).

25 More particularly, with reference to the present invention, comparisons of DNA sequences encoding the crystal toxin from distinct strains of <u>Bacillus</u> thuringiensis have revealed the existence of both conserved and variable regions. A close look to these variable regions has shown that changes are not distributed randomly over the whole gene coding for the crystal toxin, but that differences among genes

are clustered in a hypervariable region (Geiser et al. 1986).

Only few changes or no differences have been shown at the N-terminus and the C-terminus of the crystal protein genes. In fact, from the N-terminus all genes are nearly identical for approximately the first 330 aminoacids. Similarly, from about aminoacid 600 through the C-terminus the genes are largely the same.

- 10 Optimal alignment of the DNA sequences and of the deduced polypeptide sequences of these two genes and of other <u>Bacillus thuringiensis</u> genes reveals that the differences are clustered in the amino terminal halves of the molecules i.e. between aminoacid residues 280 and 640 in the case of HD-1 Dipel and HD-73 genes. This region, as hereabove mentioned, is defined as hypervariable region since it shows the maximum of variability (Geiser et al., 1986; Wabiko
- 20 On the basis of published sequences there are at least four distinct types of Lepidopteran toxins which differ substantially in this central region. Generally, the hypervariable region is, as mentioned above, in the first half of the protoxin sequence.

et al., 1986; Andrews et al., 1987).

This hypervariable region might be the result of intramolecular recombination mechanisms between very similar, but distinct genes. Moreover, the clustering of the variable subdomains in exact regions of the crystal protein toxin, strongly suggests that the exchange of hyper variable regions between genes may have caused the large variability of biological activities of different <u>Bacillus</u> thuringiensis

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crystal protein toxins. This variability is important in conferring toxic diversity and/or target insect range diversity among different Bacillus thuringiensis subspecies. The residual homology of Dipel the hypervariable region (in the case of HD-1/and HD-73 reduced to about 60% at the DNA level) should be sufficient to promote in vivo recombination.

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A similar approach has been utilized to generate, in Escherichia coli, recombinants between human

10 leukocyte interferon genes (EP 141484) and for <a href="Bacillus">Bacillus</a> alpha amylase genes (Rey et al., 1986).

Deletion analysis of Lepidopteran toxin genes has allowed the construction of much smaller proteins with full toxicity. These truncated genes show full

15 insecticidal activity only when the entire hypervariable region is present.

This analysis has been carried out for the <u>Bacillus</u> thuringiensis <u>kurstaki</u> HD-73 gene (Adang et al., 1985), the <u>Bacillus</u> thuringiensis <u>sotto</u> (Shibano et al., 1985), two genes from <u>Bacillus</u> thuringiensis

<u>kurstaki</u> HD-1 (Schnepf and Whiteley, 1985) and a Bacillus thuringiensis Berliner gene (Wabiko et al, 1986).

Since the biological activity and the action range of
the crystal toxin proteins seem to be associated to
the hypervariable region sequence, and since there is
a specific need to produce new specific <u>Bacillus</u>
thuringiensis toxin, the applicant devised, according
to the present invention, a novel way of generating
new hybrid genes and corresponding hybrid toxin
proteins, with potential different toxic specificity,
by modifying the hypervariable regions of the genes

encoding the crystal protein toxins.

Modification of the hypervariable regions of the crystal protein encoding genes could be done by site-specific mutagenesis, a technique consisting basically in introducing specific nucleotide mutations, either substitutions or deletions, in the crystal toxin hypervariable gene region, in order to

obtain aminoacid substitutions in the polypeptide

chain encoded by such mutagenized DNA sequence.

10 This technique, nevertheless, is in general very useful and efficient only if the molecular mechanism of action of the protein to be modified is known. The tridimensional structure of the protein should be known as well to be able to forecast what could be

- the consequences of the specific aminoacid substitutions or deletions in the protein to be modified. With reference to the <u>Bacillus</u> thuringiensis crystal toxin protein there are no data available as to its tridimensional structure and this
- 20 is due at least to two main reasons:
  - 1) It is quite difficult to obtain crystals of the toxin protein which can be used in a crystallographic analysis.
- 2) Last but not least, the size of the polypeptide 25 chain (135,000 dalton) would make the crystallographic analysis very laborious.

As a consequence, the site-specific mutagenesis does not seem to be very useful in providing an easy and efficient means to obtain new <u>Bacillus</u> thuringiensis

30 crystal toxins.

Another alternative approach could be a mutagenesis conducted at random on a <u>Bacillus athuringiensis</u>

crystal toxin cloned gene, but this method, lacking in specificity, does not seem to be useful to obtain new toxins characterized by insecticidal activities, since mutations introduced in the toxin gene according to mutagenesis conducted at random only affect a limited number of aminoacid residues.

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There is therefore a specific need to provide an easy and efficient method capable of producing new <a href="Bacillus thuringiensis">Bacillus thuringiensis</a> crystal toxin by modifying the

10 hypervariable region of the crystal toxin gene.

Accordingly, the present invention, by means of an in vivo-recombination process, provides a potentially unlimited number of new hybrid genes coding for new corresponding hybrid crystal protein toxins having either different insecticidal activities and/or an alterated target insect range.

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The present invention concerns new hybrid <u>Bacillus</u>
thuringiensis genes, obtained by <u>in vivo</u>
recombination, encoding new corresponding <u>Bacillus</u>
thuringiensis hybrid crystal protein toxins having
either different insecticidal acivities and/or an

30

altered insect host range.
hybrid

The new/<u>Bacillus thuringiensis</u> genes object of the present invention are obtained, as above mentioned, by means of in vivo recombination of the

- by means of <u>in vivo</u> recombination of the hypervariable region present in the <u>Bacillus</u>
- thuringiensis genes coding for the <u>Bacillus</u> thuringiensis crystal protein toxins.

The present invention also comprises new polypeptides, e.g. new hybrid crystal protein toxins

10 obtained by <u>in vivo</u> recombination of the hypervariable region of two genes coding for the <u>Bacillus</u> thuringiensis crystal protein toxins.

The two genes to be recombined in vivo can be:

- a) derived from natural <u>Bacillus</u> <u>thuringiensis</u>
- 15 strains
  - b) they can be the products of a previous  $\underline{\text{in}}$   $\underline{\text{vivo}}$  recombination event.

Another embodiment of the present invention refers to a novel process of production of new hybrid pesticidal toxins, in particular <u>Bacillus</u> thuringiensis crystal protein toxin, by <u>in vivo</u>

recombination of the hypervariable regions of genes coding for a pesticidal toxin, said genes having enough residual homology to be able to recombine <u>in</u>

<u>25</u> <u>vivo</u>.

20

30

In a preferred embodiment of this invention the two hypervariable regions to be recombined in vivo come from two different genes of <u>Bacillus thuringiensis</u> <u>kurstaki</u> and more particularly one gene is the crystal toxin encoding gene from <u>Bacillus thuringiensis kurstaki</u> HD-1 Dipel (Gene HD-1) and the other gene is the crystal toxin encoding gene from

Bacillus thuringiensis kurstaki HD-73 (Gene HD-73).

According to the present invention the in vivo recombination process applies to crystal protein encoding genes isolated from the following strains:

5 <u>Bacillus thuringiensis alesti</u>

<u>aizawai</u>

<u>canadensis</u>

dakota

darmstadiensis

10 <u>dendrolimus</u>

entomocidus

finitimus

fowleri

<u>galleriae</u>

15 indiana

israelensis

<u>Keniae</u>

kurstaki

kyushuensis

20 morrisoni

<u>ostriniae</u>

pakistani

San Diego

sotto

25 tenebrionis

thompsoni

thuringiensis

Bacillus thuringiensis tolworthi

toumanoffi

30 wuhanensi

The present invention also refers to plasmid vectors which contain two genes encoding a pesticidal protein

toxin or a part thereof, said genes having enough residual homology to be able to recombine <u>in vivo</u>. In an embodiment of the present invention these genes are the genes encoding the <u>Bacillus</u> <u>thuringiensis</u>

- 5 crystal protein toxin.
  - In a preferred embodiment of the present invention these genes are the genes encoding the <u>Bacillus</u> thuringiensis variety <u>kurstaki</u> crystal protein toxin and in a more preferred embodiment of the present
- 10 invention these genes are the genes encoding the Bacillus thuringiensis variety kurstaki HD-1 Dipel crystal protein toxin (gene HD-1) and the Bacillus thuringiensis variety kurstaki HD-73 crystal protein toxin (gene HD-73).
- These plasmid vectors containing the <u>Bacillus</u> thuringiensis DNA sequences to be recombined <u>in vivo</u> (for the reasons explained below they might be called "father plasmid vectors") represent a source of a potentially unlimited number of plasmid vectors ("son
- plasmid vectors") wherein the <u>Bacillus</u> thuringiensis

  DNA sequences have recombined <u>in vivo</u> and now these
  new hybrid DNA sequences encode new <u>Bacillus</u>

  thuringiensis hybrid crystal toxins.
- The present invention also refers to plasmid vectors

  ("son plasmid vectors" as above mentioned) containing new Bacillus thuringiensis hybrid DNA sequences, resulting from in vivo recombination of two Bacillus thuringiensis genes encoding the crystal protein toxin wherein these new hybrid DNA sequences encode

  new Bacillus thuringiensis hybrid crystal toxins
- 30 new <u>Bacillus thuringiensis</u> hybrid crystal toxins having either different insecticidal activities and/or an altered insect host range.

Objects of the present invention also are expression containing the new hybrid <u>Bacillus</u> thuringiensis DNA sequences obtained according to inventions and regulatory functions ribosome binding sites, promoters, attenuators, codons, specific SHINE-DALGARNO sequences, stop enhancers) which allow a very high expression said new hybrid Bacillus thuringiensis DNA sequences therefore a high production of their and corresponding hybrid polypeptides. 10

Expression controlling sequences useful in expressing the new hybrid <u>B. thuringiensis</u> DNA sequences of this invention include, but are not limited to, the lac system, trp system, the major operator and promoter regions of phage  $\lambda$ , the control regions of fd-coat protein, the  $\beta$ -lac system, the TAC system and other sequences known to control the expression of genes of prokaryotic or eukaryotic cells or their viruses or

15

25

30

combination thereof.

20 Another embodiment of the present invention refers to host cells transformed by plasmid vectors as here above described.

Hosts useful for preparation of the hybrid DNA sequences of the present invention by in vivo recombination include various strains of <u>E. coli</u>, <u>Pseudomonas</u>, <u>B. subtilis</u>, <u>B. thuringiensis</u>, Agrobacterium, yeasts.

For the transformation, for obtaining the hybrid Bacillus thuringiensis DNA sequences of the present invention, host cells can be recombination-proficient, recA<sup>+</sup>.

Moreover, the <u>in</u> <u>vivo</u> recombination process of

Bacillus thuringiensis DNA sequences encoding crystal toxin protein occurs in host cells rec  $A^-$ , rec  $A^-$  strains can also be used.

the use of recA strains for <u>in vivo</u> recombination 5 provides a further differentiation from the one described by Weissmann and Weber in EP-141484.

In a preferred embodiment of the present invention Escherichia coli host cells have been used.

Two father plasmid vectors called pT173 and pGEM173

10 were transformed in <u>Escherichia coli HB 101</u>, so giving rise to strains I-879 and I-878, filed at Paris Pasteur Institute on June 28, 1989. The skilled persons in the art can understand that several other host cells may be used.

15 Host cells useful for the expression of the hybrid Bacillus thuringiensis DNA sequences of the present invention are:

E. coli

Bacillus different species

20 <u>Bacillus</u> thuringiensis

Agrobacterium

Yeasts

Baculoviruses

Rhizobium

The present invention also comprises the use of the new hybrid <u>Bacillus thuringiensis</u> DNA sequences, object of this invention, for the preparation of new plasmid vectors containing these new hybrid <u>Bacillus thuringiensis</u> DNA sequences, wherein said plasmid vectors are used to transform plant cells.

New hybrid <u>Bacillus thuringiensis</u> DNA sequences of the present invention coding for new hybrid <u>Bacillus</u>

thuringiensis crystal protein toxins can in fact be engineered into plant cells to yield insect resistant plants.

The methodology for engineering plant cells is well established (see Nester E.W., Gordon M.P., Amasino R.M. and Yanofsky M.F., Ann. Rev. Plan. Physiol. 35:387-399, 1984; and EP 142 924).

Plant cells transformed by these plasmid vectors are also within the scope of the present invention.

10 The present invention comprises as well transgenic plants containing new hybrid <u>Bacillus thuringiensis</u>

DNA sequences as produced according to this invention.

In addition to be used to transform plant cells, the

15 new hybrid <u>Bacillus thuringiensis</u> DNA sequences of
the present invention can be introduced into
microorganisms capable of occupying, surviving and
proliferating in the phytosphere of plants according
to the procedures disclosed in EP-0200344.

The present invention comprises as well muteins of the hybrid <u>Bacillus thuringiensis</u> protein sequences of this invention, wherein these muteins have been obtained by standardized genetic engineering techniques (like site-specific mutagenesis, random mutagenesis, glycosilation) and their activity is reconducible to the insecticidal activity of their parent hybrid <u>Bacillus thuringiensis</u> crystal toxin proteins.

The present invention also comprises new pesticidal compositions containing new hybrid <u>Bacillus</u> thuringiensis polypeptides obtained according to this invention, in combination with suitable eccipients,

adjuvants and aggregants etc. These compositions are prepared by intimately and uniformly mixing the new hybrid <u>Bacillus</u> thuringiensis polypeptides of the present invention with suitable finely divided diluents, fillers, eccipients, disintegrating agents and the like.

As to the process of obtaining <u>in vivo</u> recombination of DNA homologous sequences we referred to a method developed by Weber and Weissman (1983) and EP-141484.

10 The methods of their invention are characterized by the steps of a) preparing a DNA fragment concatemer thereof, said fragment comprising sequence one of the parental DNA sequences from which the hybrid DNA is to be derived, an intact replicon such that the DNA fragment may be replicated in 15 host cell, and the other parental DNA sequence from which the hybrid DNA sequence is to be derived, two parental DNA sequences having sufficient homology to promote their recombination in vivo; selecting host cells that have been transformed with 20 the desired hybrid DNA sequence and isolating said hybrid DNA sequence from them. Selection of the desired host cells may be facilitated by having each parental DNA sequence associated with a different 25 antibiotic resistance marker, and growing transformed host cells on agar plates containing both antibiotics.

The hybrid DNA sequences produced by these methods clearly enable the production of novel hybrid polypeptide having a variety of uses and biological activities.

30

According to EP 141484, these methods were originally

used to produce new hybrid interferons genes by in vivo recombination of two interferon genes 

-1 and 
-2 having partial sequence homology (80%).

Thus DNA structures consisting of plasmid vector sequences flanked by the <a>-2</a> interferon gene on the one side and a portion of the -1 interferon gene on the other were transfected into E. coli host cells. Appropriate resistance markers allowed the isolation of colonies containing circular plasmides which arose 10 vivo recombination between by in the, partly homologous interferon gene sequences. In the plasmid vectors different recombinant genes were identified, all of them encoding for new hybrid interferon not accessible by traditional recombinant DNA easilv techniques. This method, according to EP 141484, 15 should be generally applicable to the formation of recombinants between not too distantly related genes. while according to the method described by Weissmann and Weber in EP-141484 host cells should be 20 recombination-proficient recA+, we have recently found that in vivo recombination of the hypervariable regions of Bacillus thuringiensis genes encoding the Bacillus thuringiensis crystal protein toxin occurs in host cells as well that are recA-. Furthermore the experimental process of the applicant thanks to the 25 use of father plasmid vector, is shorter, safer and difficult than that described by Weber less Weismann in EP-141484.

With a technique similar to that disclosed in Ep30 141484 Rey et al., 1986, obtained recombinant amylases by in vivo recombination between the genes
coding the <u>B. licheniformis</u>  $\alpha$ -amylase and the

homologous gene coding the <u>B. stearothermophilus</u>  $\alpha$  - amylase.

Another way of producing novel toxins or altering the insect host range of <u>Bacillus thuringiensis</u> toxins is provided by a method, described in EP 228838 herein enclosed as a reference, which comprises recombining in <u>vitro</u> the variable regions of two or more  $\delta$  -endotoxin genes.

Specifically exemplified in EP 228838 is the recombining of variable regions to two <u>Bacillus</u> thuringiensis <u>kurstaki</u> sequences, in particular HD-1 and HD-73, to produce chimeric <u>Bacillus</u> thuringiensis  $\delta$ -toxins with altered ranges as compared to the toxins produced by their parent DNA.

- 15 Variable regions, as used in that patent application, refers to non-homologous regions of two or more B.t.  $\delta$ -DNA sequences which upon in vitro recombination yields a DNA sequence encoding a new & -endotoxin with an altered insect host range. According to the 20 method described in EP 228838, two Bacillus thuringiensis gene showing partial homology recombined in vitro by first cutting and then religating with restriction enzymes specific Bacillus thuringiensis DNA sequences in order to obtain recombination of the non-homologous regions. 25
- However, even if this method is quite specific and allows production of new <u>Bacillus thuringiensis</u> toxins, it suffers as well of a considerable drawback since, according to an <u>in vitro</u> recombination process, only a limited amount of hybrid proteins can be produced. The method of EP 228838 requires in fact the identification of specific restriction sites in

both the genes to be recombined, while the method of the present invention, wherein recombination is carried out in vivo, allows the production of a potentially unlimited number of hybrid <u>Bacillus</u> thuringiensis toxins.

The disclosure of the present invention is intended to be read in conjunction with the references cited which are set forth in the appended bibliography.

The following examples which illustrate procedures,

- including the best mode to practice the invention, should not be considered limiting. The examples are for illustration purpose and intended to describe this invention so that it may be clearly understood.
  - The present invention provides new hybrid Bacillus
- 15 thuringiensis DNA sequences obtained by in vivo recombination of two or more different genes encoding.

  Bacillus thuringiensis crystal protein toxins.

According to the present invention, therefore, we have subcloned in plasmid vectors two truncated genes

20 of <u>Bacillus</u> <u>thuringiensis</u> variety <u>kurstaki</u>.

The source of the HD-1 Dipel gene portion was the plasmid pESAC, a derivative of pES1 (ATCC 31995) described by Schnepf and Whiteley (1981). The HD-73 gene portion derived from pJWK20 plasmid (ATCC 31997)

25 described by Kronstad and Whiteley, 1984.

pBS19 is an <u>E. coli - B. subtilis</u> shuttle vector derived from pBS42 (Wells et al., 1983; Greg Gray, unpubl.).

The gene obtained from plasmid pES1 of <u>Bacillus</u>

30 <u>thuringiensis</u> variety <u>kurstaki</u> HD-1 (gene HD-1 Dipel)

consisted in the promoter, the 5' coding sequence and the whole hypervariable coding region.

The second gene (gene HD-73) obtained from plasmid pJWK20 (Adang et al., 1985) of Bacillus thuringiensis variety <u>kurstaki</u> HD-73 was truncated in the 5' coding region and consisted, therefore, of the hypervariable region and the 3' terminal coding sequence.

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- These two genes, gene HD-1 Dipel and gene HD-73, are among those isolated from <u>Bacillus</u> thuringiensis variety kurstaki showing the greatest differences the of the nucleotide sequence of level
- 10 hypervariable region. Crystal protein toxins produced by these two different bacterial serotypes Bacillus <u>kurstaki</u> thuringiensis HD-1and Bacillus thuringiensis kurstaki HD-73 show different insecticidal activity for different species Lepidopterans. 15
  - According to the present invention the two truncated genes here above described have been cloned into plasmid pBS19 which contains a gene capable of conferring the chloramphenical resistance phenotype.
- Between the two cloned <u>Bacillus</u> thuringiensis variety 20 kurstaki genes, namely gene HD-1 Dipel and gene HDa marker gene coding for tetracycline resistance inserted. The result of this construction i. 8 plasmid vector pT173 which is shown in Figure 1.
- 25 Another plasmid, corresponding to plasmid pT173 called pGEM 173 (10.8 Kb) is shown in Figure 2. This plasmid has been obtained inserting a truncated fragment of both the <u>Bacillus</u> thuringiensis <u>kurstaki</u> HD-1 gene and the Bacillus thuringiensis kurstaki HD-30 73 gene in the plasmid vector pGEM4Z (Promega, WI, USA Plasmid pT173, once introduced in

Madison,

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Escherichia coli cells synthetizes a polypeptide of 65

Kd which is recognized by specific antibodies for the Bacillus thuringiensis crystal toxin protein. This polypeptide has been identified as the product of Bacillus thuringiensis variety kurstaki 'HD-1, which, as here above described, is truncated downstream of the hypervariable region. The partial homology between the two hypervariable regions of gene HD-1

Diepel and gene HD-73 (at nucleotide sequence level

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this homology is 62,2%) should be sufficient to be 10 able to promote an <u>in vivo</u> recombination process of the two truncated genes.

It is clear that since this is an <u>in vivo</u> recombination process, plasmid vectors like plasmid pT173 can produce an unlimited number of different

15 recombinant hybrid <u>Bacillus</u> <u>thuringiensis</u> DNA sequences encoding for their corresponding hybrid <u>Bacillus</u> <u>thuringiensis</u> polypeptides having either different insecticidal activity and/or an altered target insect range.

10

20 Plasmid pT173, therefore, has been transformed in suitable Escherichia coli host cell.

In one embodiment of the present invention, these  $\underline{Escherichia}$   $\underline{coli}$  host cells are recombination proficient rec  $A^+$ , but according to some more recent

25 results, <u>in vivo</u> recombination of <u>Bacfllus</u> thuringiensis DNA sequences occurs also in cells which are rec A<sup>-</sup>.

The recombination deficient rec A cells (instead of recombination proficient rec A+) have the advantage that the hybrid DNA is not rearranged. In this way accuracy of recombination is obtained to the DNA level, avoiding casual rearrangment and then the

occurence of unexpected and unwanted sequences.

After many generations plasmid DNA is extracted and digested with the restriction enzyme Nru I.

Since recombinant plasmid carrying new <u>Bacillus</u>

<u>5 thuringiensis</u> hybrid DNA sequences obtained upon <u>in</u>

<u>vivo</u> recombination have lost the DNA restriction site

recognized by Nru I (as it will be explained in a

more detailed way further on), treatment with the

endonuclease Nru I allows selection of plasmids which

10 have recombined in vivo.

These recombinant plasmids, being the only ones still circular after the Nru I treatment, will be able successively to transform <u>E. coli</u> cells producing their colonies which will be resistant to

15 chloramphenicol.

The same procedure applies to plasmid pGEM 173, with the only difference that <u>E. coli</u> transformants are selected for resistance to ampicillin instead of chloramphenicol.

20 According to the method hereabove described, we succeeded in isolating many different recombinants whose hybrid DNA sequences are reported in Figure 6 a - e.

#### MATERIALS AND METHODS

### 25 BACTERIAL STRAINS

The following bacterial strains and their rec A derivative were used for transformation:

### Bacterial strains

Escherichia coli strains were: HB101 (F- hsdS20

30 recA13 ara-14 proA2 leuB6 lacY1 galK2 rpsL20 xy1-5 mt1 -1 sup E44)

JM103: (lac proAE)  $\triangle$  (lac pro), thi, strA, supE,

endA, sbeB, hsdR-, F'traD36, proAB, lac19, Z \( \trace{AM15} \)

294 endA thi pro hsdR hsdM hsm

294recA endA thi pro hsdR hsdM hsm recA

Escherichia coli host cells were made competent and transformed according to Hanahan 1985.

#### CULTURE MEDIA

LB medium (per liter: Difco Bacto-tryptone 10g; Difco Bacto-yeast extract 5g; NaCl 5g)

For growth of strains with plasmid pT173,

10 tetracycline 12.5 /ug/ml or chloramphenicol 10 /ug/ml were added to LB medium.

For growth of strains with plasmid pGEM-173 ampicillin was used at 100 µg/ml.

#### PLASMID DNA EXTRACTION

15 Plasmid DNA preparation has been done by the alkaline lysis method adapted to larger samples and followed, for sequencing, by a PEG precipitation (Birnboim and Doly, 1979).

## RESTRICTION ENZYMES, DIGESTION, ELECTROPHORESIS

20 Restriction enzymes were from Boehringer Mannheim GmbH II (FRG) and from Bethesda Research Laboratories (Maryland, USA).

Restriction enzymes digestion, ligation and other treatments during plasmid construction have been done

25 following the suggested protocols of the supplier.

Restriction patterns generated from the digestion of plasmid DNA were resolved on 0.7% agarose gels (agarose from Bethesda Research Laboratories). DNA samples were electrophoresed in TBE buffer (0.3M 30 Tris-borate pH 8.3, 2 mM EDTA) and stained with

30 Tris-borate pH 8.3, 2 mM EDTA) and stained with ethidium bromide. When necessary electrophoresis was performed on acrylamide gels (6%) in TBE buffer.

## DNA SEQUENCE ANALYSIS

Fragments of DNA, originated from recombinant plasmids derived from pT173 and pGEM-173, obtained by MindITI digestion. A band corresponding to DNA of approximately 2.9 Kb was 5 separated agarose gel electrophoresis, recovered by electroelution and subsequently digested with EcoRV. A band of DNA of approximately 700 bp was purified by acrylamide gel electrophoresis (6%) electroeluted and 10 ligated to the plasmid pGEM-4Z (Promega, Madison, W1, USA) digested with **SmaI** and treated with phosphatase. Sequencing was performed by the chain termination method of Sanger et al. (1977) adapted to plasmid DNA (Chen and Seeburg, 1985).

15 Sequenase (United States Biochemical) is known, in the dideoxi-chain elongation reaction.

#### IMMUNOBLOTTING

with the antiserum.

25

The method of Towbin et al. (1974) was used to detect the crystal protein immunologically.

- 20 Proteins resolved by Sodium dodecyl sulphatepolyacrylamide gel electrophoresis were transferred electrophoretically to nitrocellulose sheets washed with 50 mM Tris hydrochloride-200 mM NaCl containing 0.1% Nonidet P-40 and then incubated
  - After a wash with the same buffer, the nitrocellulose sheets were incubated with peroxidase-conjugated sheep anti-rabbit immunologlobulin G antiserum (United States Biochemical Co., Cleveland, Ohio).
- 30 The immuno-complexes were then visualized in the presence of hydrogen peroxide and 4-chloro-1-naphtol as substrates (GIBCO Laboratories, Grand Island

N.Y., Bethesda Research Laboratories, Inc. Gaithersburg, Md.)

## PLASMID CONSTRUCTION

pT is a pBS19 derivative: a 1424 bp <u>Eco</u> RI - <u>Ava</u> I (with even ends in <u>Ava</u> I) fragment of pBR322 bearing tetracycline resistance was inserted in pBs 19 digested with <u>Eco</u> RI and <u>Sac</u> I (the latter digestions followed by treatments able to obtain even ends).

pT73 is a pT derivative obtained by inserting in the

10 Eco RI site of pT an Eco RI fragment of approximately 5400 bp obtained from pJWK20 (Fig. 1). The fragment comprises the last two thirds of the HD73 toxin gene starting from residue 1383 of the sequenced region (Adang et al., 1985). pT1 was obtained as follows: pT

was digested to completion with <u>Sma</u> I, partially with <u>Bam</u> HI and ligated to a fragment of approximately 1900 bp derived from pESAC and corresponding to the first portion of HD-1 Dipel toxin gene from nucleotide residue 291 to residue 2215 of the sequenced region (Schnepf et al., 1985).

The fragment was obtained by digestion of pESAC with Hind III followed by treatment with Klenow to make it blunted and then by restriction with Bam HI.

pT173, the plasmid used for the <u>in vivo</u> construction of recombinant genes, derived from the insertion of the 2500bp <u>Bam</u> HI-Bam HI fragment of pT1, comprising the first part of the HD-1 Dipel gene and the last two thirds of the resistance tetracycline gene, in pT73 completely cleaved with <u>Bam</u> HI and dephosphorilated with pancreatic phosphatase to avoid

re-insertion of the original Bam HI fragment.

TetR and CmR transformants obtained in E. coli 294

racA strain were examined by plasmid extraction and restriction analysis.

Plasmids pT1, pT73 and pT173 were not able to direct the synthesis of a complete endotoxine polypeptide but only of a 65-68 kdal fragment which immunoreacted with antibodies raised against the pure toxic crystal in pT1 and pT173.

The tetracycline resistant gene was inserted in inverted orientation to ensure that the HD73

10 truncated sequence could not be expressed from external expression control regions in pT73 and pT173.

### GENERATION OF HYBRID GENES

The sequences of HD-1 Dipel and HD73 genes in pT173 share a region of homology, i.e. the last 696 bp of HD-1 Dipel sequence and the first 707 bp of the HD-73 sequence. Between these partial direct repeats there is the tetracycline resistance determinant characterized by the unique Nru I site.

- 20 Figure 3 shows the protocol followed to generate the recombinant plasmids with the hybrid toxin genes.

  The pT173 plasmid was introduced in a recombination proficient background by transformation of <u>E. coli</u>
  294 competent cells.
- A single colony Tet<sup>R</sup> and Cm<sup>R</sup> has been inoculated in LB (supplemented with chloramphenicol) and grown for about 40 generations. Plasmid DNA extracted from the cells was digested with Nru I. In this way the molecules not subjected recombination and having the intact tetracycline gene were linearized. Only circular molecules could replicate and transform the

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294 recA competent cells to CmR.

This allows selection of plasmids which have recombined in vivo: they have lost the restriction site recognized by the endonuclease Nru I and according to that they remain circular.

- The CmR transformants were screened for sensitivity to tetracycline with the aim of obtaining recombinants between the partially homologous regions of the truncated toxin genes (boxed in Figure 3). In this way we expected to reconstitute an entire hybrid
- of the HD-1 Dipel gene and two thirds from the carboxyterminal region of the HD-73 gene. The hypervariable region was expected to be a different hybrid region for each clone able to express a
- 15 polypeptide immunoreacting with specific antibodies.

  We isolated 13 Cm<sup>R</sup> Tet<sup>S</sup> clones, examined them for the production of a polypeptide immunoreacting with polyclonal antibodies raised against HD-73 toxic crystals and for the presence of recombinant plasmids 20 (pTHy).

## ANALYSIS OF THE RECOMBINANT OBTAINED

Escherichia coli cells transformed with the recombinant plasmids have been analyzed by immunoblotting to identify all the clones capable of

- 25 synthesizing a polypeptide chain of 135 kd having the immunological properties of the <u>Bacillus</u> thuringiensis crystal toxin protein. The method of Towbin et al (1974) was used to detect the crystal protein immunologically.
- 30 According to this method 10 positive clones were identified.

In order to characterize these positive clones

identified, recombinant plasmids have been purified and their DNA was digested with restriction enzymes in order to obtain their restriction maps. Plasmid purification and restriction enzyme digestion have been done according to the methods indicated previously in description of the present patent application.

All the plasmids analyzed were identified as recombinant plasmids obtained by  $\underline{\text{in}}$   $\underline{\text{vivo}}$ 

10 recombination of the hypervariable regions of the two <a href="Bacillus thuringiensis kurstaki"><u>Bacillus thuringiensis kurstaki</u></a> genes contained either in plasmid pT173 or pGEM173.

These two "father plasmids" are in fact sources of a potentially unlimited number of new hybrid <u>Bacillus</u>

15 thuringiensis DNA sequences obtained by in vivo recombination.

All these recombinant plasmids were further characterized by determining the nucleotide sequence of DNA fragments corresponding to the hypervariable regions wherein, according to the present invention,

DNA sequence analysis has been done according to the method indicated in the description of this patent

application.

recombination has occurred.

20

In figure 4 the HD-1 and the HD-73 <u>Bacillus</u> thuringiensis DNA sequences are aligned to maximize the matches (represented by a vertical line). The regions of cross-over are indicated by a box.

While most of the recombinant hybrid DNA sequences

30 obtained and identified are different from each
other, some of them were completely identical and we
can not exclude that they might be "brothers"

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resulting from a unique recombination process.

As indicated in Figure 4, the <u>in vivo</u> recombination process has occurred through all the hypervariable region. So far, of all the recombinant hybrid <u>Bacillus</u> thuringiensis DNA sequences analyzed, excluding those which recombined in the same region, 10 hybrids used a different region of cross-over. The DNA sequences of these 10 new recombinant hybrid <u>Bacillus</u> thuringiensis genes are reported in Figure 6

10 a - e.

20

These recombinant hybrid sequences have been called respectively: HY45, HY3, HY6, HY53, HY21, HY32, HY2, HY 127, HY 126, HY 5.

The aminoacid sequence of the proteins coded by these 15 recombinant hybrid <u>Bacillus thuringiensis</u> genes can be easily deduced from their nucleotide sequence.

A comparison of the deduced aminoacid sequences is reported in Figure 5 which represents the optimal alignment of these polypeptide products deduced from the DNA sequence of their parental recombinant hybrid genes. According to the specific site where the recombination process has occurred, the hybrid protein corresponds in the hyper variable region

either to the protein encoded by the gene HD-1 Dipcl

25 or to the protein encoded by the gene HD 73.

Two cases have been identified, HY6, HY64 and HY 127, HY 21, where, while the hybrid recombinant genes have a different nucleotide sequence, the polypeptide chains are identical. At the aminoacid level, therefore, the real number of new recombinants

30 therefore, the real number of new recombinants encoding hybrid <u>Bacillus</u> thuringiensis crystal protein toxins is eight. These hybrid genes encode

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proteins which are different from their hybrid parental natural Bacillus thuringiensis crystal toxin from any and other known Bacillus thuringiensis crystal toxin protein.

- Accordingly, these new hybrid proteins might have either different insecticidal activities and/or altered insect host range as compared to the natural crystal toxin proteins produced bv Bacillus thuringiensis variety kurstaki HD-1 Dipel or Bacillus thuringiensis variety kurstaki HD-73.
- The experimental procedure described in the present invention in order to generate hybrid recombinants between different Bacillus thuringiensis genes can be applied to other couples of <a href="Bacillus">Bacillus</a> thuringiensis

10

- 15 genes or different recombinant products, to generating new polypeptides possibly having spectra of toxic activity different from that of the parental products. Among the 8 new different hybrid Bacillus thuringiensis crystal protein toxins · isolated,
- preliminary results indicate that some of them have a 20 specific toxic activity against Ostrinia nubilalis larvae (European corn borer).

It is clear to those skilled in the art that direct sequence analysis of other recombinants combined with assays of toxicity against different Lepidopteran 25 targets could lead to the identification of new toxins with either a different insecticidal activity or an altered insect host range. Those skilled in the art will appreciate, therefore, that the invention described herein and the methods of practising it 30 specifically described are susceptible of variations modifications and other than as specifically

described.

It is to be understood that the invention includes all these variations and modifications which are intended to be fully within the scope of the following claims.

The numerals set forth below represent figure numbers for the appended drawings.

Fig. 1 - Shows the construction of pT173 plasmid.

10 PT is a derivative of pBS19 able to express chloramphenicol resistance (Cm) and tetracycline resistance (Tet, heavy black arrow) in <u>E. coli</u> and <u>B. subtilis</u>.

The open box is the HD-73 toxin coding sequence starting from residue 1383. The direction of transcription is indicated. The heavy line represents the HD-73 sequences downstream the toxin gene.

Dashed box represents the HD-1 Dipel sequence starting from residue 291 to residue 2215. The

20 direction of transcription is indicated. The interrupted arrows indicate the region of partial homology.

Fig. 2 - Shows the restriction map of plasmid pGEM173 (10.8 kb)

- 25 This plasmid is obtained by inserting a truncated fragment of the HD-73 gene, more particularly an EcoRI-ecoRI fragment derived from plasmid pJWK20 (as for the construction of plasmid pT173) in the plasmid vector pGEM 4Z Promega.
- 30 The truncated HD-1 gene inserted in plasmid pGEM-173 is a BamHI-BamHI obtained from plasmid pT173. This BamHI-BamHI fragment only includes part of the TctR

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gene, which contains a NrvI restriction site.

Fig. 3 - Is a schematic representation of the process of  $\underline{\text{in}}$   $\underline{\text{vivo}}$  recombination of the present invention.

- 5 Plasmid PT 173 (13 kb) carrying the <u>Bacillus</u> thuringiensis HD-1 and HD-73 truncated sequences, separated by the marker gene for the resistance to tetracycline, has been introduced by transformation in <u>E. coli</u> (RecA+) cells wherein <u>in vivo</u>
- 10 recombination of the two <u>Bacillus thuringiensis</u> DNA sequences occurs.

PTHY represents isolated recombinant vectors carrying genes coding for a new hybrid <u>Bacillus</u> thuringiensis toxin.

15 The heavy closed boxes represent the partially homologous regions, open for HD-73, dashed for HD-1 Dipel.

20

Fig. 4 - Shows the alignment of the two <u>Bacillus</u> thuringiensis DNA sequences HD-1 and HD-73 in order to give the greatest homology.

Vertical dashed lines indicate matches; boxes represent, for each recombinant obtained by the present method, regions wherein the recombination has occurred; numbers in bold above boxes represent the

- 25 new recombinant obtained HY 6, HY 15, HY 107, HY 53.

  DNA sequence nucleotides are indicated and numbered according to the original numeration given in the published DNA sequences HD-1 (Schnepf et al., 1985) and HD-73 (Adang et al., 1985).
- 30 Fig. 5 Shows the optimal alignment of polypeptide deduced from the sequences of HD-1 Dipel and HD-73 genes.

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Panel A (modified after Geiser et al., In 1986) schematically represented the alignment of the entire polypeptides; a vertical line represents an unmatched residue, short bars near the horizontal lines deletions.

Panel B represents the aminoacid sequences aligned for the region corresponding to the partial homology region, at the DNA sequence level, used recombination in pT173 and pGEM173.

The sequence indicated in this figure corresponds to 10 the region underlined in Panel A.

An aminoacid residue identical in HD-1 Dipel and HD-73 or in the hybrid products is denoted by a bar.

- represents an aminoacid residue deleted to obtain 15 maximum alignment.

Figs 6 a-e Show the DNA nucleotide sequences of hypervariable regions of hybrid Bacillus thuringiensis genes obtained by in vivo recombination according to the present invention.

The DNA sequence has been determined sperimentally on 20 both DNA strands for all the recombinant hybrid genes obtained.

HD-1 DNA sequence is in bold letters. Numeration is as follows: Position 1 corresponds to nucleotide 1521

25 of the HD-1 DNA sequence (Schnepf, 1985) while the last nucleotide corresponds to nucleotide 2091 of the HD-73 DNA sequence (Adang et al., 1985).

#### CLAIMS

- 1. Hybrid DNA sequences characterized in that they are obtained by <u>in vivo</u> recombination of two genes coding for an insecticidal toxin protein.
- 5 Hybrid DNA sequences according to Claim wherein said hybrid DNA sequences have been obtained by in vivo recombination of two Bacillus thuringiensis genes coding for the Bacillus thuringiensis crystal toxin protein.
- 3. Hybrid DNA sequences according to Claims 1-2, wherein said hybrid DNA sequences have been obtained by in vivo recombination of two <u>Bacillus</u> thuringiensis variety <u>kurstaki</u> genes coding for the <u>Bacillus</u> thuringiensis crystal toxin protein.
- 4. Hybrid DNA sequences according to Claims 1-3, wherein said DNA sequences have been obtained by in vivo recombination of the Bacillus thuringiensis variety kurstaki HD-1 Dipel gene and the Bacillus thuringiensis var. kurstaki HD-73 gene, both genes coding for the Bacillus thuringiensis crystal toxin protein.
- 5. Hybrid DNA sequences according to Claims 1-4 wherein said hybrid DNA sequences have been obtained by in vivo recombination of the hypervariable regions of the <u>Bacillus thuringiensis</u> var. <u>kurstaki HD-1</u> Dipel gene and of <u>Bacillus thuringiensis</u> var. <u>kurstaki HD-73</u> gene, both genes coding for the <u>Bacillus thuringiensis</u> crystal toxin protein.
- 6. Hybrid DNA sequence HY 5 according to Claims
  30 1-5 characterized by an hypervariable region having the following nucleotide sequence:

LOCUS

HY5

713 BP

ENTERED 5/22/87

ORIGIN

IN VIVO RECOMBINANT HD1/HD73 #5

- 1 AATTCBCATT CCCTTTATTT BBGAATBCBG BBAATBCABC TCCACCCBTA CTTBTCTCAT

  61 TAACTBBTTT BGGBATTTTT ABAACATTAT CTTCACCTTT ATATABAABA ATTATACTTB

  121 BTTCABGCCC AAATAATCAB BAACTBTTTB TCCTTBATBG AACBBABTTT TCTTTTBCCT

  181 CCCTAACGAC CAACTTBCCT TCCACTATAT ATAGACAAAG BGGTACAGTC BATTCACTAG

  5 241 ATBTAATACC BCCACABAAT AACAACGTGC CACCTAGGCA AGGATTTAGT CATCGATTAA

  301 GCCATGTTCC AATGTTTCGT TCAGGCTTTA GTAATAGTAG TGTAAGTATA ATAAGAGCTC

  361 CTATGTTCTC TTGGATACAT CGTAGTGCTG AATTTAATAA TATAATTGCA TCGGATAGTA

  421 TTACTCAAAT CCCTGCAGTG AAGGGAAACT TTCTTTTTAA TGGTTCTGTA ATTCAGGAC

  481 GAGGGTATAT TGGAGGGGC TTAGTTAGAT TAAATAGTAG TGGAAATAAC ATTCAGAATA,

  841 GAGGGTATAT TGAAGTTCCA ATTCACTTCC CATCGACATC TACCAGATAT CGAGTTCGTG

  461 TTTCCAATAC AGTACCAGCT ACAGCTACGT CATTAGATAA TCTACAATCA AGT
  - 7. Hybrid DNA sequence HY 45 according to Claims 1-5 characterized by an hypervariable region having the following sequence:

15 Locus

HY45

707 BP

ENTERED 5/23/89

ORIGIN

IN VIVO RECOMBINANTS HD-1/HD73 #45,66

- 1 AATTCBCATT CCCTTTATTT BOGAATBCBG BBAATBCABC TCCACCCBTA CTTBTCTCAT

  61 TAACTBGTTT BOGGATTTTT AGAACATTAT CTTCACCTTT ATATAGAABA ATTATACTTB

  121 BTTCABGCCC AAATAATCAB BAACTBTTTB TCCTTBATBG AACBGABTTT TCTTTTBCCT

  181 CCCTAACBAC CAACTTBCCT TCCACTATAT ATAGACAAAB BBGTACABTC BATTCACTAB

  241 ATBTAATACC BCCACABGAT AATABTBTAC CACCTCBTBC BBGATTTAGC CATCBATTBA

  301 BTCATBTTAC AATBCTGAGC CAAGCAGCTB BAGCAGTTTA CACCTTBAGA BCTCCAACGT

  361 TTTCTTBGAT ACATCGTAGT GCTGAATTTA ATAATATAAT TGCATCGGAT AGTATTACTC

  421 AAATCCCTGC AGTGAAGGGA AACTTTCTTT TTAATGGTTC TGTAATTTCA GGACCAGGAT

  481 TTACTGGTGG GGACTTAGTT AGATTAAATA GTAGTGGAAA TAACATTCAG AATAGAGGGT

  541 ATATTGAAGT TCCAATTCAC TTCCCATCGA CATCTACCAG ATATCGAGTT CGTGTACGGT

  401 ATGCTTCTGT AACCCCGATT CACCTCAACG TTAATTGGGG TAATTCATCC ATTTTTTCCA

  461 ATACAGTACC AGCTACAGCT ACGTCATTAG ATAATCTACA ATCAAGT
  - 8. Hybrid DNA sequence HY 3 according to Claims 1-5 characterized by an hypervariable region having the following nucleotide sequence:

LOCUS HY3 710BP ENTERED 5/23/89

ORIGIN IN VIVO RECOMBINANTS HD-1/HD73 #3,4,7

- 1 AATTCBCATT CCCTTTATTT BBBAATBCBB BBAATBCABC TCCACCCBTA CTTBTCTCAT
  61 TAACTBGTTT BBGBATTTTT ABAACATTAT CTTCACCTTT ATATABAAGA ATTATACTTB
  121 BTTCABGCCC AAATAATCAB BAACTBTTB TCCTTBATBB AACBGABTTT TCTTTTBCCT
  181 CCCTAACBAC CAACTTBCCT TCCACTATAT ATABACAAAB BBGTACABTC BATTCACTAB
  241 ATBTAATACC BCCACABGAT AATABTBTAC CACCTCBTBC BBGATTTABC CATCBATTBA
  301 BTCATBTTAC AATBCTBABC CAAGCAGCTB BADCABTTTA CACCTTBABA BCTCCAACBT
  361 TTTCTTBGCA BCATCBCABT BCTBAATTTA ATAATATAAT TCCTTCATCA CAAATTACAC
  421 AAATACCTTT AACAAAATCT ACTAATCTTB BCTCTBBAAC TTCTBTCGTT AAAGGACCAG
  481 BATTTACTGG TGGGGACTTA GTTAGATTAA ATAGTAGTGG AAATAACATT CAGAATAGAG
  601 GGTATGCTTC TGTAACCCCG ATTCACCTCA ACGTTAATTG GGGTAATTCA TCCATTTTTT
- 9. Hybrid DNA sequence HY 21 according to Claims
  1-5 characterized by an hypervariable region having
  the following nucleotide sequence:

661 CCANTACAGT ACCAGCTACA GCTACGTCAT TAGATAATCT ACAATCAAGT

- 15 LOCUS HY21 713BP ENTERED 5/23/89
  ORIGIN IN VIVO RECOMBINANT HD-1/HD73 #21
- 1 AATTCBCATT CCCTTATTT BGBAATBCBB BBAATBCABC TCCACCCBTA CTTBTCTCAT
  61 TAACTGGTTT BGGGATTTTT AGAACATTAT CTTCACCTTT ATATABAABA ATTATACTTB
  121 BTTCABBCCC AAATAATCAB BAACTBTTTB TCCTTBATBB AACBBABTTT TCTTTTBCCT
  181 CCCTAACBAC CAACTTBCCT TCCACTATAT ATAGACAAAB BGGTACABTC BATTCACTAB
  20 241 ATBTAATACC BCCACABGAT AATABTBTAC CACCTCBTBC BBGATTTABC CATCBATTBA
  301 BCCATGTTTC AATGTTTCGT TCAGGCTTTA GTAATAGTAG TGTAAGTATA ATAAATAGAGCTC
  361 CTATGTTCTC TTGGATACAT CGTAGTGCTG AATTTAATAA TATAATTGCA TCGGATAGTA
  421 TTACTCAAAT CCCTGCAGTG AAGGGAAACT TTCTTTTTAA TGGTTCTGTA ATTTCAGGAC
  481 CAGGATTTAC TGGTGGGGAC TTAGTTAGAT TAAATAGTAG TGGAAATAAC ATTCAGAATA
  541 GAGGGTATAT TGAAGTTCCA ATTCACTTCC CATCGACATC TACCAGATAT CGAGTTCGTG
  601 TACGGTATGC TTCTGTAACC CCGATTCACC TCAACGTTAA TTGGGGTAAT TCATCCATTT
  - 10. Hybrid DNA sequence HY 32 according to Claims 1-5 characterized by an hypervariable region having the following nucleotide sequence:

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LOCUS HY32 707BP ENTERED 5/23/89
ORIGIN IN VIVO RECOMBINANT HD-1/HD73 #32

11. Hybrid DNA sequence HY 6 according to Claims 1-5 characterized by an hypervariable region having the following nucleotide sequence:

15 LOCUS HT6 710BP UPDATED 5/23/89
ORIGIN IN VIVO RECOMBINANT HD-1/HD73 #6

1 AATTCBCATT CCCTTATTT BOGAATBCOO BARATGCAGC TCCACAACAA CGTATTGTTG
61 CTCAACTAGG TCAGGGCGTG TATAGAACAT TATCGTCCAC TTTATATAGA AGACCTTTA
121 ATATAGGGAT ARATAATCAA CAACTATCTG TTCTTGACGG GACAGAATTT GCTTATGGAA
181 CCTCCTCAAA TTTGCCATCC GCTGTATACA GAAAAAGCGG AACGGTAGAT TCGCTGGATG
241 AAATACCGCC ACAGAATAAC AACGTGCCAC CTAGGCAAGG ATTTAGTCAT CGATTAAGCC
301 ATGTTTCAAT GTTTCGTTCA GGCTTTAGTA ATAGTAGTGT AAGTATAATA AGAGCTCCTA
361 TGTTCTCTTG GATACATCGT AGTGCTGAAT TTAATAATAT AATTGCATCG GATAGTATTA
421 CTCAAATCCC TGCAGTGAAG GGAAACTTTC TTTTTAATGG TTCTGTAATT TCAGGACCAG
481 GATTTACTGG TGGGGACTTA GTTAGATTAA ATAGTAGTGG AAATAACATT CAGAATAGAG
641 GGTATGCTTC TGTAACCCCG ATTCACCTCA ACGTTAATTG GGGTAATTCA TCCATTTTT
661 CCAATACAGT ACCAGCTACA GCTACGTCAT TAGATAATCT ACAATCAAGT

12. Hybrid DNA sequence HY 53 according to Claims 1-5 characterized by an hypervariable region having the following nucleotide sequence:

25

20

LOCUS HY53 710BP UPDATED 5/22/89
ORIGIN IN VIVO RECOMBINANTS HD-1/HD73

#53,64,107

1 AATTCOCATT CCCTTTATTT BOGAATOCOG BOGATOCADC TCCACAACAA CGTATTGTTG
61 CTCAACTAGG TCAGGGCGTG TATAGAACAT TATCGTCCAC TTTATATAGA AGACCTTTTA
121 ATATAGGGAT AAATAATCAA CAACTATCTG TTCTTGACGG GACAGAATTT GCTTATGGAA
181 CCTCCTCAAA TTTGCCATCC GCTGTATACA GAAAAAGCQG AACGGTAGAT TCGCTGGATG
241 AAATACCGCC ACAGAATAAC AACGTGCCAC CTAGGCAAGG ATTTAGTCAT CGATTAAGCC
301 ATGTTCAAT GTTTCGTTCA GGCTTTAGTA ATAGTAGTGT AAGTATAATA AGAGCTCCTA
361 TGTTCTCTTG GATACATCGT AGTGCTGAAT TTAATAATAT AATTGCATCG GATAGTATTA
421 CTCAAAATCCC TGCAGTGAAG GGAAACTTTC TTTTTAATGG TTCTGTAATT TCAGGACCAG
481 GATTTACTGG TGGGGACTTA GTTAGATTAA ATAGTAGTGG AAATTACATT CAGAATAGAG
541 GGTATATTGA AGTTCCAATT CACTTCCCAT CGACATCTAC CAGATATCGA GTTCGTGTAC
601 GGTATGCTTC TGTAACCCCG ATTCACCTCA ACGTTAATTG GGGTAATTCA TCCATTTTTT
661 CCAATACAGT ACCAGCTACA GCTACGTCAT TAGATAACTT ACAATCAAGT

13. Hybrid DNA sequence HY 2 according to Claims
1-5 characterized by an hypervariable region having
15 the following nucleotide sequence:

LOCUS HY2 713BP UPDATED 5/23/89
ORIGIN IN VIVO RECOMBINANT HD-1/HD73 #2

1 AATTCBCATT CCCTTTATTT BOBAATBCBB BBAATBCABC TCCACCCBTA CTTBTCTCAT
61 TAACTBGTTT BOBGATTTTT AGAACATTAT CTTCACCTTT ATATABAABA ATTATACTTB
121 BTTCABBCCC AAATAATCAB BAACTBTTTB TCCTTBATBB AACBBABTTT TCATTTBCCT
181 CCCTAACBAC CAACTTBCCT TCCACTATAT ATABACAAAB BGGTACABTC BATTCACTAB
241 ATBTAATACC BCCACAGBAT AATAGTBTGC CACCTAGGCA AGGATTTAGT CATCGATTAA
301 GCCATGTTC AATGTTTCGT TCAGGCTTTA GTAATAGTAG TGTAAGTATA ATAAGAGCTC
361 CTATGTTCTC TTGGATACAT CGTAGTGCTG AATTTAATAA TATAATTGCA TCGGATAGTA
421 TTACTCAAAT CCCTGCAGTG AAGGGAAACT TTCTTTTTAA TGGTTCTGTA ATTTCAGGAC
481 CAGGATTTAC TGGTGGGGAC TTAGTTAGAT TAAATAGTAG TGGAAATAAC ATTCAGAATA
541 GAGGGTATAT TGAAGTTCCA ATTCACTTCC CATCGACATC TACCAGATAT CGAGTTCGTG
601 TACGGTATGC TCCTGTAACC CCGATTCACC TCAACGTTAA TTGGGGTAAT TCATCCATTT

14. Hybrid DNA sequence HY 127 according to Claims 1-5 characterized by an hypervariable region having the following nucleotide sequence:

WO 91/01087 PCT/EP90/01145

LOCUS HY127 713BP ENTERED 5/23/89

ORIGIN IN VIVO RECOMBINANT HD-1/HD73 #127

1 AATTEBCATT CCCTTTATTT BOGAATBCBB BBAATBCABC TCCACCCBTA CTTBTCTCAT
61 TAACTBGTTT BOGGATTTTT AGAACATTAT CTTCACCTTT ATATAGAABA ATTATACTTB
121 BTTCABBCCC AAATAATCAB BAACTBTTTB TCCTTBATBB AACBBABTTT TCTTTTBCCT
181 CCCTAACBAC CAACTTBCCT TCCACTATAT ATAGACAAAB BOGTACAGTC BATTCACTAB
241 ATBTAATACC BCCACAGBAT AATAGTBTAC CACCTCBTBC BOGATTTABT CATCGATTAA
301 GCCATGTTCC AATGTTTCGT TCAGGCTTTA GTAATAGTAG TGTAAGTATA ATAAGAGCTC
361 CTATGTTCTC TTGGATACAT CGTAGTGCTG AATTTAATAA TATAATTGCA TCGGATAGTA
421 TTACTCAAAT CCCTGCAGTG AAGGGAAACT TTCTTTTTAA TGGTTCTGTA ATTTCAGGAC
481 CAGGATTTAC TGGTGGGGAC TTAGTTAGAT TAAATAGTAG TGGAAATAAC ATTCAGAATA
541 GAGGGTATAT TGAAGTTCCA ATTCACTTCC CATCGACATC TACCAGATAT CGAGTTCGTG
601 TACCGGTATGC TTCTGTAACC CCGATTCACC TCAACGTTAA TTGGGGTAAT TCATCCATTT
641 TTTCCAATAC AGTACCAGCT ACAGCTACGT CATTAGATAA TCTACAATCA AGT

15. Hybrid DNA sequence HY 126 according to Claims 1-5 characterized by an hypervariable region having the following nucleotide sequence:

15 LOCUS HY126 707 BP ENTERED 5/22/89

#### ORIGIN IN VIVO RECOMBINANT HD-1/HD 73 #126

1 AATTCBCATT CCCTTTATTT BOBAATBCBB BBAATBCABC TCCACCCBTA CTTBTCTCAT
61 TAACTBGTTT BGBGATTTTT ABAACATTAT CTTCACCTTT ATATABAAGA CCTTTTAATA
121 TAGGGATAAA TAATCAACAA CTATCTGTC TTGACGGGAC AGAATTTGCT TATGGAACCT
181 CCTCAAATTT GCCATCCGCT GTATACAGAA AAAGCGGAAC GGTAGATTCG CTGGATGAAA
241 TACCGCCACA GAATAACAAC GTGCCACCTA GGCAAGGATT TAGTCATCGA TTAAGCCATG
301 TTTCAATGTT TCGTTCAGGC TTTAGTAATA GTAGTGTAAG TATAATAAGA GCTCCTATGT
361 TCTCTTGGAT ACATCGTAGT GCTGAATTTA ATAATAAAT TGCATCGGAT AGTATTACTC
421 AAATCCCTGC AGTGAAGGGA AACTTTCTT TTAATGGTC TGTAATTTCA GGACCAGGAT
481 TTACTGGTGG GGACTTAGTT AGATTAAATA GTAGTGGAAA TAACATTCAG AATAGAGGGT
541 ATATTGAAGT TCCAATTCAC TTCCCATCGA CATCTACCAG ATATCGAGTT CGTGTACGGT
661 ATACAGTACC AGCTACAGCT ACGTCATTAG ATAATCTACA ATCAAGT

- 16. Hybrid insecticidal crystal toxin protein encoded by the hybrid DNA sequences according to Claims 1-15.
- 30 17. Hybrid insecticidal crystal toxin proteins having substantially the immunological properties of the hybrid insecticidal crystal toxin protein

10

30

according to Claim 16.

- 18. Muteins of the hybrid insecticidal crystal toxin proteins according either to Claim 16 or Claim 17 wherein said muteins have been obtained by standardized genetic engineering techniques such as site-specific mutagenesis, random mutagenesis, site-specific glycosilation, and their activities are reconducible to the insecticidal activities of their parental hybrid insecticidal crystal toxin proteins according to either Claim 16 or Claim 17.
- 19. Plasmid vectors characterized in that they contain two genes coding for an insecticidal toxin protein and having enough residual homology to be able to recombine in vivo, wherein said genes upon in vivo recombination produce the hybrid DNA sequences of Claims 1-15.
- 20. Plasmid vectors according to Claim 19 further characterized in that the genes coding for the insecticidal toxin protein are accociated and 20 separated on said plasmid vectors by a DNA fragment acting as an antibiotic resistance marker.
- 21. Plasmid vectors according to Claim 19 wherein the genes coding for an insecticidal toxin protein are <u>Bacillus thuringiensis</u> genes coding for the <u>Bacillus thuringiensis</u> crystal toxin protein.
  - 22. Plasmid vectors according to Claim 21 wherein the <u>Bacillus thuringiensis</u> genes are <u>Bacillus thuringiensis</u> variety <u>kurstaki</u> genes coding for the <u>Bacillus thuringiensis</u> variety <u>kurstaki</u> crystal toxin protein.
  - 23. Plasmid vectors according to Claim 22 wherein the <u>Bacillus thuringiensis</u> variety <u>kurstaki</u>

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genes have been derived from <u>Bacillus</u> <u>thuringiensis</u> variety <u>kurstaki</u> HD-1 Dipel and from <u>Bacillus</u> <u>thuringiensis</u> variety <u>kurstaki</u> HD 73, said genes coding both for crystal toxin protein.

- 5 24. Plasmid vectors according to Claim 23 wherein said plasmid vectors are pT173 and pGEM173.
- 25. Plasmid vector characterized in that it contains a first gene coding for a first insecticidal toxin protein and a second gene coding for a second insecticidal toxin protein, having enough amology in order to be able to recombine in vivo, so that it produces hybrid DNA sequences coding for a third insecticidal toxin protein, wherein said first gene is characteristic of the hypervariable region of a first bacterial strain, while the said second gene is characteristic of the hypervariable region of a second bacterial strain substantially different from the said first strain.
- 26. Vector according to claim 25 characterized 20 in that said first bacterial strain and said second bacterial strain belong to the same genera, preferably <u>Bacillus</u> type.
- 27. Vector according to claim 26 characterized in that said first bacterial strain and said second 25 bacterial strain belong to the same species, preferably <u>Bacillus thuringiensis</u> type.
  - 28. Vector according to claim 27 characterized in that said first bacterial strain and said second bacterial strain belong to the same subspecies, preferably of the <u>Bacillus thuringiensis kurstaki</u> type.

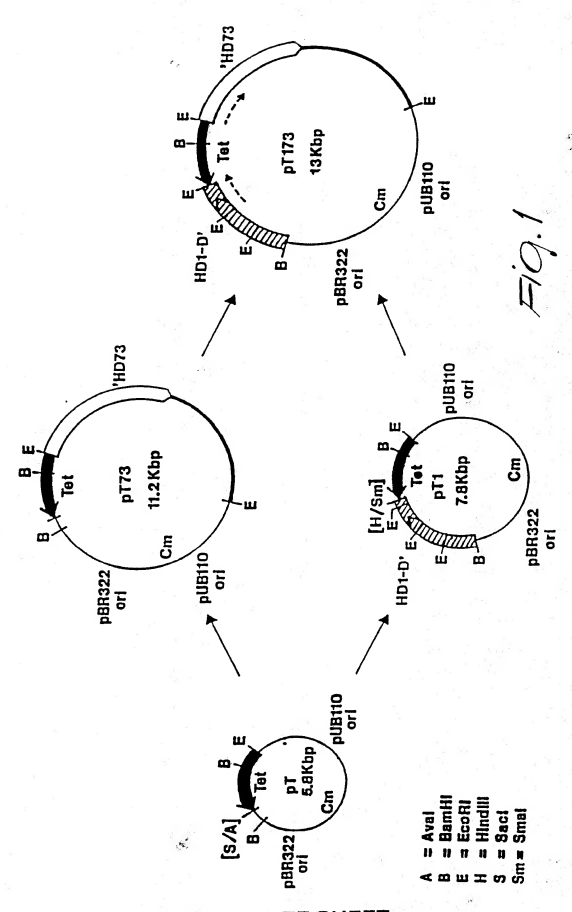
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29. Plasmid expression vectors characterized in

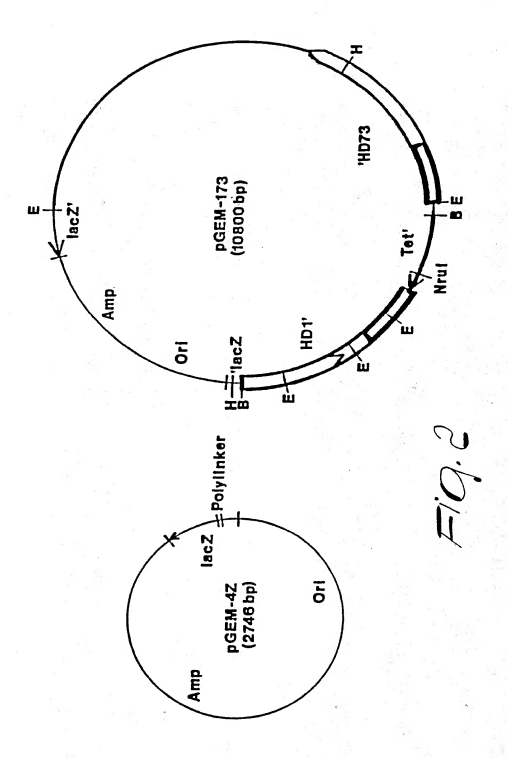
that they have been obtained by <u>in vivo</u> recombination of the two genes coding for an insecticidal toxin protein contained in plasmid vectors according to Claims 19 and 25 wherein said plasmid vectors obtained by <u>in vivo</u> recombination contain the hybrid DNA sequences of Claims 1-15.

- 30. Plasmid expression vectors containing the hybrid DNA sequences according to Claims 1-15 under the control of expression regulatory functions.
- 31. Plasmid expression vectors according to Claim 29 wherein said expression regulatory functions include, but are not limited to, the lac system, the Trp system, the major operator and promoter regions of phage A, the tac system, the B-lac system.
- 32. Plasmid expression vectors containing the hybrid DNA sequences according to Claims 1-15 wherein said plasmid expression vectors are used to transform plant cells.
- 33. Plant cells transformed by plasmid 20 expression vectors according to Claim 29.
  - 34. Transgenic insect resistant plants containing the hybrid DNA sequences according to Claims 1-15.
- 35. Genetically engineered plant colonizing 25 microorganism containing the hybrid DNA sequences according to Claim 1-15.
  - 36. Host cells transformed by the plasmid expression vectors according to Claim 29.
- 37. Host cells according to Claim 36 wherein 30 said host cells are various strains of <u>Bacillus</u>, including <u>B. subtilis</u>, <u>B. thuringiensis</u>, yeasts, <u>Agrobacterium</u>, baculoviruses, <u>Rhizobium</u>.

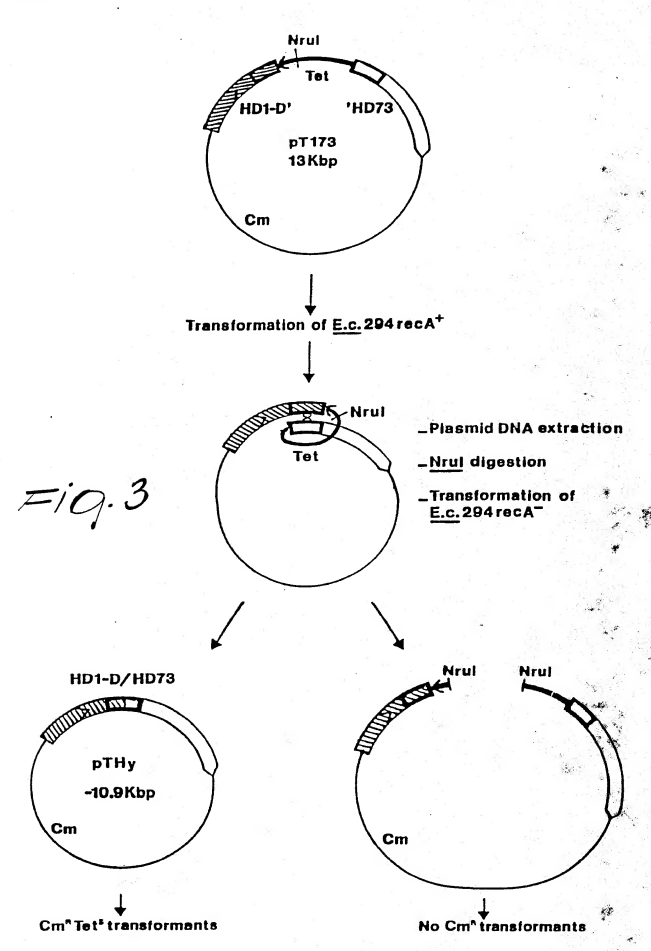
- 38. Host cells according to Claim 36 wherein said host cells are Escherichia Coli host cells.
- 39. Host cells according to Claim 38 wherein said host cells can be recombinant unproficient cells  $5 \text{ rec } A^-$ .
  - 40. <u>Escherichia</u> <u>coli</u> host cells transformed with plasmid vector pT173.
  - 41. Escherichia coli host cells transformed with plasmid vector pGEM 173.
- 10 42. A process for preparing the hybrid DNA sequences of Claims 1-15 by <u>in vivo</u> recombination of two or more genes coding for an insecticidal toxin protein and having enough residual homology to be able to recombine <u>in vivo</u>.
- 15 '43. A process according to Claim 42, wherein said genes coding for an insecticidal protein are <a href="Bacillus thuringiensis">Bacillus thuringiensis</a> genes coding for the <a href="Bacillus thuringiensis">Bacillus thuringiensis</a> crystal toxin protein.
- 44. Pesticidal compositions and formulations
  20 containing the hybrid crystal toxin proteins of
  Claims 16 and 17 in combination with suitable
  eccipients, diluents, fillers, aggregant and the
  like.
- 45. Use of the hybrid crystal toxin proteins of 25 Claim 17 to control and to combat insect pest.



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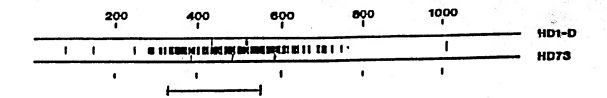


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HD 1 Dipe HD 73 gen	
	6.15 64.53.107
1521	AATTCGCATTCCCTTTATTTGGGAATGCGGGGAATGCAGCTCCACCCGTACTTGT CTC
1383	ATTCACTTTTCCGCTATATGGAACTATGGGAAATGCAGCTCCAGAACAACGTATTGTTG
1579	ATTAACT GGTTTGGGGATTTTTAGAACATTATCTTCACCTTTATATAGAAGAATTATAC
1442	CTCAACTAGGTCAGGGCGTGTATAGAACATTATCGTCCACTTTATATAGAAGACCT T
1638	TTGGTTCAGGCCCAAATAATCAGGAACTGTTTGTCCTTGATGGAACGGAGTTTTCTTTTG
1499	TTAATATAGGGATAAATAATCAACAACTATCTGTTCTTGACGGGACAGAATTTGCTTATG
1698	CCTCCCTAACGACCAACTTGCCTTCCACTATATATAGACAAAGGGGTACAGTCGATTCAC
1559	GAACCT CCTCAAATTTGCCATCCGCTGTATACAGAAAAAGCGGAACGGTAGATTCGC
1758	TAGATGTAATACCGCCACAGATAATAGTGTACCACCTCGTGCCGGATTTAGCCATCGAT
1616	TGGATGAATACCGCCACAGAATAACAACGIGCCACCTAGGCAAGGATTTAGTCATCGAT
	21
1818	TGAGTCATGTTACAATGCTGAG CCAAGC AGCTGGAGCAGTTTA CACCTTGAGAG
1676	TAAGCCATGTTTCAATGTTTCGTTCAGGCTTTAGTAATAGTAGTGTAAGTATAATAAGAG
1872	CTCCAACGTTTCCTTGGCAGCATCGCAGTGCTGAATTTAATAATATAATTCCTTCATCAC
1736	CTCCTATGTTCTTGCATACATCGTAGTGCTGAATTTAATAATAATTGCATCGGATA
1932	AAATTACACAAATACCTTTAACAAAATCTACTAATCTTGGCTCTGGAACTTCTGTCGTTA
1796	GTATTACTCAAATCCCTGCAGTGAAGGGAAACTTTCTTTTTAATGG TTCTGTAATTT
1992	AAGGACCAGGATTTACAGGAGGAGATATTCTTCGAAGAACTTCACCTGGCCAGA TTTC
1853	daggaccaggatttadreggeggacttagttagattagatagtagtagaaataacattc
2050	AACCTTAAGAGTAAAT A TT ACTGCA CCAT TATCACAAAGATATCGGG
1913	AGAATAGAGGGTATATTGAAGTTCCAATTCACTTCCCATCGACATCTACCAGATATCGAG
	TAAGAATTCGCTACGCTTCTACTACAAATTTACAATTCCATACATCAATTGACGGAA G
2097	
1973	TTCGTGTACGGTATGCTTCTGTAACCCCGATTCACCT CA ACGTTAATTGGGGTAATTC
2155	ACCTATTAATCAGGGTAATTTTTCAGCAAC TATGAGTAGTGGGAGTAATTTACAGTCCG
2031	ATCCATTTTTCCAATACAGTACCAGCTACAGCTACGTCATTAGA TAATCTACAATCAA
2214	GA
2090	GT
Matches = Length =	
	=10.4



E

# HYBRID PROTEINS FROM RECOMBINANT GENES

HD1-D HY7,3,4 HY66,45 HY32 HY127,21 HY2 HY5 HY126 HY6,53,64,	107	FAFPLFBNAGNAAPPV~LVSLTBLBIFRTLSSPLYRRIILBSGPNNG	Q-sAYG
HD73		-TY-TMQQRI-AQL-Q-VYTP^FNI-I	***
HD1-D HY7,3,4 HY66,45 HY32- HY127,21	<b>3</b> 93	TTNLPSTIYRQRGTVDSLDVIPPQDNSVPPRAGFSHRLSHVTM^LS	GPSNSS-SI 11
HY5 HY126 HY6,53,64,	107		GF8NSS-SI I 1 GF8NSS-8I I 1
D73  -   D1-D (	<b>4</b> 52	SWOMENSAEFNINI I PESQITQI PLTKETNLESGTSVVK BPSFT GGDIL	
HY7,3,4 HY66,45 HY32 HY127,21 HY2 HY5 HY126			n-ru-s-huidh-d n-ru-s-huidh-d n-ru-s-huidh-d n-ru-s-huidh-d n-ru-s-huidh-d n-ru-s-huidh-d
1Y7,3,4 1Y66,45 1Y32 1Y127,21 1Y2 1Y5		ITAPLEQRYRV^RIRYASTINLQFHTSIDERPINGENF^EATHSEES -BY-IHFPSTBT-Y-VRYRYASYTPIHLNYNNG-851-8HTVPÄTA* -BY-IHFPSTBT-Y-VRYRYASYTPIHLNYNNG-851-8HTVPÄTA* -BY-IHFPSTBT-Y-VRYRYASYTPIHLNYNNG-851-8HTVPÄTA* -BY-IHFPSTBT-Y-VRYRYASYTPIHLNYNNG-851-8HTVPÄTA* -BY-IHFPSTBT-Y-VRYRYASYTPIHLNYNNG-851-8HTVPÄTA* -EY-IHFPSTST-Y-VRYRYASYTPIHLNYNNG-851-8HTVPÄTA* -EY-IHFPSTST-Y-VRYRYASYTPIHLNYNNG-851-8HTVPÄTA* -EY-IHFPSTST-Y-VRYRYASYTPIHLNYNNG-851-8HTVPÄTA* -EY-IHFPSTST-Y-VRYRYASYTPIHLNYNNG-851-8HTVPÄTA*	L2-DNFG L2-DNFG L2-DNFG L2-DNFG L2-DNFG L2-DNFG L2-DNFG L2-DNFG L3-DNFG

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LOCUS HY45 707 BP 5/23/89 ENTERED ORIGIN IN VIVO RECOMBINANTS HD-1/HD73 #45.66 1 AATTCBCATT CCCTTTATTT BBBAATBCBB BBAATBCABC TCCACCCBTA CTTBTCTCAT 61 TAACTEGTTT EGGEATTTTT AGAACATTAT CTTCACCTTT ATATAGAAGA ATTATACTTE 121 STTCABBCCC AAATAATCAB BAACTETTTB TCCTTBATEB AACBBABTTT TCTTTTBCCT 181 CCCTAACGAC CAACTTBCCT TCCACTATAT ATAGACAAAG GGGTACAGTC GATTCACTAG 241 ATGTAATACC BCCACABGAT AATAGTGTAC CACCTCGTGC GGGATTTAGC CATCGATTGA 301 BTCATETTAC AATECTBABC CAABCABCTB BAGCABTTTA CACCTTBABA BCTCCAACGT 421 AAATCCCTGC AGTGAAGGGA AACTTTCTTT TTAATGGTTC TGTAATTTCA GGACCAGGAT 481 TTACTGGTGG GGACTTAGTT AGATTAAATA GTAGTGGAAA TAACATTCAG AATAGAGGGT 541 ATATTGAAGT TCCAATTCAC TTCCCATCGA CATCTACCAG ATATCGAGTT CGTGTACGGT 601 ATGCTTCTGT AACCCCGATT CACCTCAACG TTAATTGGGG TAATTCATCC ATTTTTTCCA 661 ATACAGTACC AGCTACAGCT ACGTCATTAG ATAATCTACA ATCAAGT

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ORIGIN

IN VIVO RECOMBINANTS HD-1/HD73 #3,4,7

1 AATTCBCATT CCCTTTATTT BEBAATBCBB BBAATBCABC TCCACCCBTA CTTBTCTCAT
61 TAACTBBTTT BBBGATTTTT ABAACATTAT CTTCACCTTT ATATABAAGA ATTATACTTB
121 BTTCABBCCC AAATAATCAB BAACTBTTTB TCCTTBATBB AACBBABTTT TCTTTTBCCT
181 CCCTAACBAC CAACTTBCCT TCCACTATAT ATABACAAAB BBBATTTABC CATCBATTBA
241 ATBTAATACC BECACABBAT AATABTBTAC CACCTCBTBC BBBATTTABC CATCBATTBA
301 BTCATBTTAC AATBCTBABC CAABCABCTB BABCABTTTA CACCTTBABA BCTCCAACBT
361 TTTCTTBBCA BEATCBCABT BCTBAATTTA ATAATATAAT TCCTTCATCA CAAATTACAC
421 AAATACCTTT AACAAAATCT ACTAATCTTB BCTCTBBAAC TTCTBTCBTT AAABBACCAB
481 BATTTACTGG TGGGGACTTA GTTAGATTAA ATAGTAGTGG AAATAACATT CAGAATAGAG
541 GGTATATTGA AGTTCCAATT CACTTCCCAT CGACATCTAC CAGATATCGA GTTCGTGTAC
601 GGTATGCTTC TGTAACCCCG ATTCACCTCA ACGTTAATTG GGGTAATTCA TCCATTTTTT
661 CCAATACAGT ACCAGCTACA GCTACGTCAT TAGATAATCT ACAATCAAGT

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F19.6a

LOCUS HY6 710 BP UPDATED 5/23/89 ORIGIN IN VIVO RECOMBINANT HD-1/HD73 #6 1 AATTCBCATT CCCTTTATTT BBBAATBCBB BAAATGCAGC TCCACAACAA CGTATTGTTG 61 CTCAACTAGG TCAGGGCGTG TATAGAACAT TATCGTCCAC TTTATATAGA AGACCTTTTA 121 ATATAGGGAT AAATAATCAA CAACTATCTG TTCTTGACGG GACAGAATTT GCTTATGGAA 181 CCTCCTCAAA TTTGCCATCC GCTGTATACA GAAAAAGCGG AACGGTAGAT TCGCTGGATG 241 AAATACCGCC ACAGAATAAC AACGTGCCAC CTAGGCAAGG ATTTAGTCAT CGATTAAGCC 301 ATGTTTCAAT GTTTCGTTCA GGCTTTAGTA ATAGTAGTGT AAGTATAATA AGAGCTCCTA 361 TGTTCTCTTG GATACATCGT AGTGCTGAAT TTAATAATAT AATTGCATCG GATAGTATTA 421 CTCAAATCCC TGCAGTGAAG GGAAACTTTC TTTTTAATGG TTCTGTAATT TCAGGACCAG 481 GATTTACTGG TGGGGACTTA GTTAGATTAA ATAGTAGTGG AAATAACATT CAGAATAGAG 541 GGTATATTGA AGTTCCAATT CACTTCCCAT CGACATCTAC CAGATATCGA GTTCGTGTAC 601 GGTATGCTTC TGTAACCCCG ATTCACCTCA ACGTTAATTG GGGTAATTCA TCCATTTTTT 661 CCAATACAGT ACCAGCTACA GCTACGTCAT TAGATAATCT ACAATCAAGT

LOCUS HY53

710 BP

UPDATED 5/22/89

ORIGIN

IN VIVO RECOMBINANTS HD-1/HD73 #53,64,107

AATTCBCATT CCCTTTATTT BBBAATBCBB BBAATBCABC TCCACAACAA CGTATTGTTG
61 CTCAACTAGG TCAGGGCGTG TATAGAACAT TATCGTCCAC TTTATATAGA AGACCTTTTA
121 ATATAGGGAT AAATAATCAA CAACTATCTG TTCTTGACGG GACAGAATTT GCTTATGGAA
181 CCTCCTCAAA TTTGCCATCC GCTGTATACA GAAAAAGCGG AACGGTAGAT TCGCTGGATG
241 AAATACCGCC ACAGAATAAC AACGTGCCAC CTAGGCAAGG ATTTAGTCAT CGATTAAGCC
301 ATGTTCAAT GTTTCGTTCA GGCTTTAGTA ATAGTAGTGT AAGTATAATA AGAGCTCCTA
361 TGTTCTCTTG GATACATCGT AGTGCTGAAT TTAATAATAT AATTGCATCG GATAGTATTA
421 CTCAAATCCC TGCAGTGAAG GGAAACTTC TTTTTAATGG TTCTGTAATT TCAGGACCAG
481 GATTTACTGG TGGGGACTTA GTTAGATTAA ATAGTAGTGG AAATAACATT CAGAATAGAG
541 GGTATATTGA AGTTCCAATT CACTTCCCAT CGACATCTAC CAGATATCGA GTTCGTGTAC
601 GGTATGCTC TGTAACCCCG ATTCACCTCA ACGTTAATTG GGGTAATTCA TCCATTTTTT
661 CCAATACAGT ACCAGCTACA GCTACGTCAT TAGATAATCT ACAATCAAGT

Fig. 66

LOCUS HY21 713 BP ENTERED 5/23/89 ORIGIN IN VIVO RECOMBINANT HD-1/HD73 #21 1 AATTCGCATT CCCTTTATTT BBBAATBCBB BBAATBCABC TCCACCCBTA CTTGTCTCAT 61 TAACTEGITT GEGGATTITT AGAACATTAT CTTCACCTTT ATATAGAGA ATTATACTTE 121 STTCASSCCC AAATAATCAS SAACTSTTTS TCCTTSATSS AACSSASTTT TCTTTTSCCT 181 CCCTAACBAC CAACTTBCCT TCCACTATAT ATABACAAAB BBBTACABTC BATTCACTAB 241 ATBTAATACC BCCACABGAT AATABTBTAC CACCTCSTBC BBGATTTABC CATCBATTBA 301 GCCATGTTTC AATGTTTCGT TCAGGCTTTA GTAATAGTAG TGTAAGTATA ATAAGAGCTC 361 CTATGTTCTC TTGGATACAT CGTAGTGCTG AATTTAATAA TATAATTGCA TCGGATAGTA 421 TTACTCAAAT CCCTGCAGTG AAGGGAAACT TTCTTTTTAA TGGTTCTGTA ATTTCAGGAC 481 CAGGATTTAC TGGTGGGGAC TTAGTTAGAT TAAATAGTAG TGGAAATAAC ATTCAGAATA 541 GAGGGTATAT TGAAGTTCCA ATTCACTTCC CATCGACATC TACCAGATAT CGAGTTCGTG 601 TACGGTATGC TTCTGTAACC CCGATTCACC TCAACGTTAA TTGGGGTAAT TCATCCATTT 661 TTTCCAATAC AGTACCAGCT ACAGCTACGT CATTAGATAA TCTACAATCA AGT 11

LOCUS

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ORIGIN

IN VIVO RECOMBINANT HD-1/HD73 #32

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Locus	HY 126	707 E	P		ENTERED	5/22/89
ORIGIN	IN VIVO	RECOMBINANT	HD-1/HD73	#126		
1	AATTCBCATT	CCCTTTATTT	BBBAATBCBB	BBAATBCABC	TECACCEBTA	CTTBTCTCAT
						CCTTTTAATA
						TATGGAACCT
						CTGGATGAAA
						TTAAGCCATG
						GCTCCTATGT
						AGTATTACTC
						GGACCAGGAT
		GGACTTAGTT				
541	ATATTGAAGT	TCCAATTCAC	TTCCCATCGA	CATCTACCAG	ATATCGAGTT	CGTGTACGGT
		AACCCCGATT				
661	ATACAGTACC	AGCTACAGCT	ACGTCATTAG	ATAATCTACA	ATCABGT	ALLITICOA
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LOCUS

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ORIGIN

IN VIVO RECOMBINANT HD1/HD73 #5

1 AATTCBCATT CCCTTTATTT BBBAATBCBB BBAATBCABC TCCACCCBTA CTTBTCTCAT
61 TAACTBBTTT BBBBATTTTT ABAACATTAT CTTCACCTTT ATATABAABA ATTATACTTB
121 BTTCABBCCC AAATAATCAB BAACTBTTTB TCCTTBATBB AACBBABTTT TCTTTTBCCT
181 CCCTAACBAC CAACTTBCCT TCCACTATAT ATABACAAAB BBBTACABTC BATTCACTAB
241 ATBTAATACC BCCACABAAT AACAACGTGC CACCTAGGCA AGGATTTAGT CATCGATTAA
301 GCCATGTTC AATGTTTCGT TCAGGCTTTA GTAATAGTAG TGTAAGTATA ATAAGAGCTC
361 CTATGTTCTC TTGGATACAT CGTAGTGCTG AATTTAATAA TATAATTGCA TCGGATAGTA
421 TTACTCAAAT CCCTGCAGTG AAGGGAAACT TTCTTTTTAA TGGTTCTGTA ATTTCAGGAC
481 CAGGATTTAC TGGTGGGGAC TTAGTTAGAT TAAATAGTAG TGGAAATAAC ATTCAGAATA
541 GAGGGTATAT TGAAGTTCCA ATTCACTTCC CATCGACATC TACCAGATAT CGAGTTCGTG
601 TACGGTATGC TTCTGTAACC CCGATTCACC TCAACGTTAA TTGGGGTAAT TCATCCATTT
661 TTTCCAATAC AGTACCAGCT ACAGCTACGT CATTAGATAA TCTACAATCA AGT

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F19.6d

LOCUS

HY127

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LOCUS	HY2	713 B	P		UPDATED !	5/23/89
ORIGIN	OVIV NI	RECOMBINANT	HD-1/HD73	#2		
ř <b>1</b>	AATTCBCATT	CCCTTTATTT	868AATBC88	BBAATBCABC	TCCACCCBT	CTTBTCTCAT
61	TAACTEGTTT	BEGGATTTTT	ABAACATTAT	CTTCACCTTT	ATATABAABA	ATTATACTTE
121	BTTCABBCCC	AAATAATCAB	BAACTETTTB	TCCTTBATEB	AACBBAGTT	TCTTTTBCCT
181	CCCTAACBAC	CAACTTBCCT	TCCACTATAT	ATABACAAAB	BESTACAST	BATTCACTAB
241	ATBTAATACC	<b>BCCACAGBAT</b>	AATAGTGTGC	CACCTAGGCA	AGGATTTAGT	CATCGATTAA
301	GCCATGTTTC	AATGTTTCGT	TCAGGCTTTA	GTARTAGTAG	TGTAAGTAT	ATAAGAGCTC
361	CTATGTTCTC	TTGGATACAT	CGTAGTGCTG	AATTTAATAA	TATAATTGC	ATEGERATAGE
421	TTACTCAAAT	CCCTGCAGTG	AAGGGAAACT	TTCTTTTTAA	TGGTTCTGT	ATTTCAGGAC
481	CAGGATTTAC	TGGTGGGGAC	TTAGTTAGAT	DATDATAAAT	TGGAAATAAC	ATTCAGAATA
541	GAGGGTATAT	TGAAGTTCCA	ATTCACTTCC	CATCGACATC	TACCAGATAT	CGAGTTCGTĞ
601	TACGGTATGC	TTCTGTAACC	CCGATTCACC	TCAACGTTAA	TTGGGGTAAT	TCATCCATTT
661	TTTCCAATAC	AGTACCAGCT	ACAGCTACGT	CATTAGATAA	TCTACAATCA	TDA

ORIGIN IN VIVO RECOMBINANT HD-1/HD73 #127

1 AATTCBCATT CCCTTTATTT BBBAATBCBB BBAATBCABC TCCACCCBTA CTTETCTAT

61 TAACTBETTT BGBBATTTTT ABAACATTAT CTTCACCTTT ATATABAABA ATTATACTTB

121 BTTCABBCCC AAATAATCAB BAACTBTTB TCCTTBATBB AACBBABTTT TCTTTTBCCT

181 CCCTAACBAC CAACTTBCCT TCCACTATAT ATABACAAAB BGBTACABTC BATTCACTAB

241 ATBTAATACC BCCACABBAT AATABTBTAC CACCTCBTBC BGBATTTABT CATCGATTAA

301 GCCATGTTC AATGTTCGT TCAGGCTTTA GTAATAGTAG TGTAAGTATA ATAAGAGCTC

361 CTATGTTCTC TTGGATACAT CGTAGTGCTG AATTTAATAA TATAATTGCA TCGGATAGTA

421 TTACTCAAAT CCCTGCAGTG AAGGGAAACT TTCTTTTTAA TGGTTCTGTA ATTCAGGAC

481 CAGGATTTAC TGGTGGGGAC TTAGTTAGAT TAAATAGTAG TGGAAATAAC ATTCAGAATA

541 GAGGGTATAT TGAAGTTCCA ATTCACTTCC CATCGACATC TACCAGATAT CGAGTTCGTG

**601 TACGGTATGC TTCTGTAACC CCGATTCACC TCAACGTTAA TTGGGGTAAT TCATCCATTT** 

661 TITCCARTAC AGTACCAGCT ACAGCTACGT CATTAGATAA TCTACAATCA AGT

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Fig. 6e

	JECT MATTER (if several classifica		
According to International Pate Int.Cl. 5	nt Classification (IPC) or to hoth Natio A01N63/02; C12N15		5/62
II. FIELDS SEARCHED			
	Minimum Do	ocumentation Scarched <sup>7</sup>	
Classification System		Classification Symbols	5- <b>46</b> 75.
Int.Cl. 5	C07K ; C12N		
	Documentation Searched of the Extent that such Documents	other than Minimum Ducumentation ents are Included in the Fields Searched <sup>8</sup>	.d.
III. DOCUMENTS CONSIDERE			
Category ° Citation of D	ocument, 11 with indication, where appr	ropriate, of the relevant passages 12	Relevant to Claim No. 13
Y EP,A,14 see page 1, 4	1484 (BIOGEN N.V.) e 8, line 18 - page 1	l1, line 10; figures	1-5, 18-19, 21-23,
see page (cited	e 12, line 22 - page in the application)	13, line 12	42-45
see page see page	B838 (MYCOGEN CORP.) 2 2, lines 15 - 22; c 3 7; examples 3-5 3 in the application)	laims	1-5, 18-19, 21-23, 25-39 42-45
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"I." document which may throw which is cited to establish the citation or other special reast of document referring to an or other means	eral state of the art which is not ar relevance hed on or after the international doubts on priority claim(s) or he publication date of another son (as specified) ral disclosure, use, exhibition or the international filing date but	or priority date and not in conflict with the cited to understand the principle or theory invention  "X" document of particular relevance; the claim cannot be considered novel or cannot be considered novel or cannot be considered to involve an inventive step  "V" document of particular relevance; the claim cannot be considered to involve an inventive document is combined with one or more cited.	ment published after the international filing date date and not in conflict with the application but nderstand the principle or theory underlying the of particular relevance; the claimed invention considered novel or cannot be considered to involve an inventive step when the is combined with one or particular relevance; the claimed invention considered to involve an inventive step when the is combined with one or more other such documents combination being obvious to a person skilled member of the same patent family  3 0. 10, 30  [ Authorized Officer
ate of the Actual Completion of the	International Search	I Day of Marie	
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ternational Searching Authority EUROPEAN	N PATENT OFFICE	Signature of Authorized Officer ANDRES S.M.	3

III. DOCUME	NTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)	
Category o	Citation of Document, with indication, where appropriate, of the relevant passages	Relevant to Claim No
Y	PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF USA. vol. 86, June 1989, WASHINGTON US pages 4037 - 4041; GE, A.Z. et al.: "Location of the Bombyx mori specificity domain on a Bacillus thuringiensis delta-endotoxin protein" see the whole document	1-5, 18-19, 21-23, 25-39
		42-45
Y	NUCLEIC ACIDS RESEARCH.  vol. 11, no. 16, 1983, ARLINGTON, VIRGINIA  pages 5661 - 5669; WEBER,H. & WEISSMANN,C.:  "Formation of genes coding for hybrid proteins  by recombination between related, cloned genes  in E.coli"  see the whole document	1-5, 18-19, 21-23, 25-39
	(cited in the application)	42-45
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# ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

EP 9001145 SA 38941

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on

The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

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EP-A-228838	15-07-87	JP-A-	62143689	26-06-87	
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